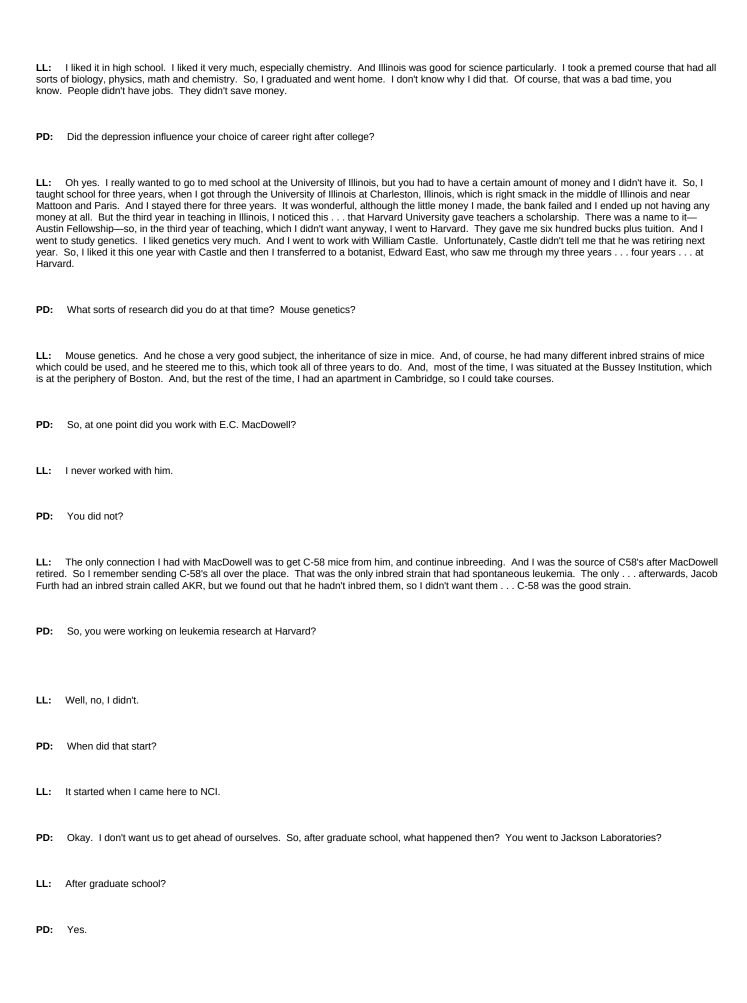
Law, Lloyd 2001

Dr. Lloyd Law Oral History 2001

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Natio	nal Cancer Institute Oral History Interview Project
Inter	view with Lloyd Law
Cond	ucted on January 30, 2001, by Peggy Dillon
In Be	thesda, MD
PD:	Good afternoon Dr. Law.
LL:	Good afternoon. I'm happy to have you here.
	Thank you. I'm happy to be here. Today I would like to talk with you about your work at the National Cancer Institute and your contributions to er research. And I would like to start by asking you about your background before you joined the National Cancer Institute in 1947. Could you tell mabout your upbringing in Pennsylvania and your education at the University of Illinois and Harvard?
	I was born in Ford City, Pennsylvania, which is western Pennsylvania, on the Allegheny River near Pittsburgh. And after where I was born, a the first month, my mother decided she wanted to live in California. So, we went to California.
PD:	When you were a month old?
we ca	Yes. And I went we stayed there until I got through kindergarten. And the reason we came back is when I was six or seven the reason ame back was my father sustained a bad burn, so we came back despite the objections by my mother. She didn't want to come back to western sylvania.
PD:	You returned to Ford City?
LL:	Ford City.
PD:	Okay.
didn't was the Illinois Cham	Because my father was a railroader and he wanted to come back here and do the railroad, which he did. He got a job at the Pittsburgh Plate Glass cany and we started to school, the elementary school there. So, in a two-room schoolhouse, grades one to four and grades five to eight. And we have running water at that time, which wasn't unusual. So, I graduated from high school and, fortunately, one of the people I got to know very well he coach of the track team. And he was thirty years old. He never graduated from high school, but somehow he got admitted to the University of s. So he invited me to apply and I applied, seventy-five dollars tuition a semester. And when the time came, we hopped the train for Urbanangarian or than, except the trip to California, I had never been that far. Illinois was about 400 miles from Ford City. So, I got and got a job at a dollar an hour, working in the football training room and graduated from the University of Illinois. I think, in 1931.

PD: How did you decide to study science?

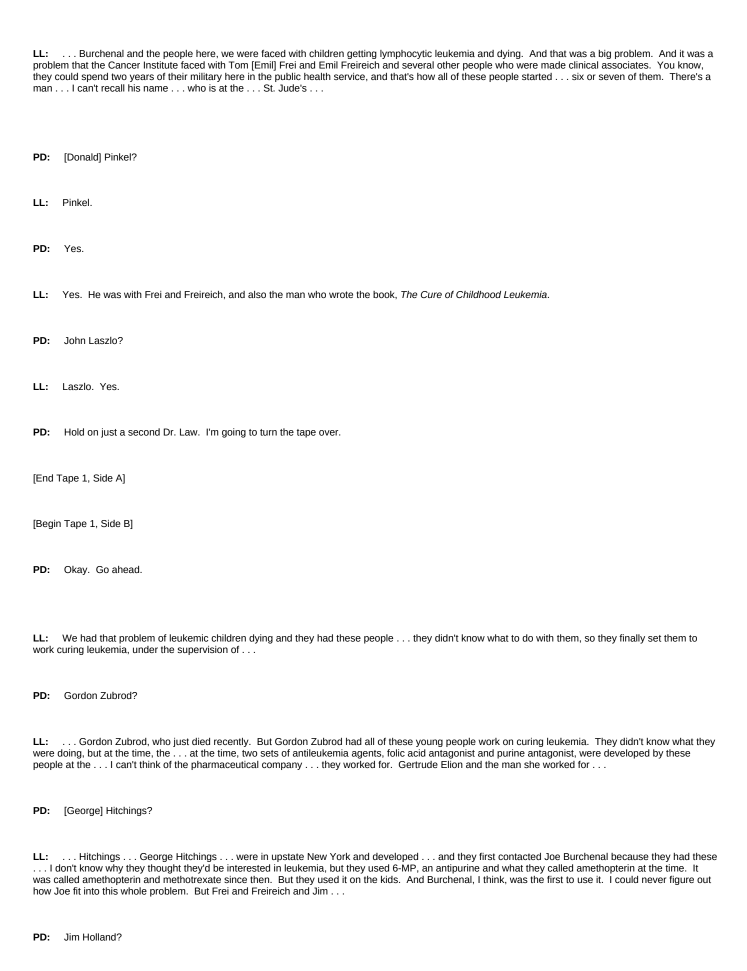


LL: Well, I, you know, they were looking for Army buddies, and I had a low number number eight, in Bar Harbor. No, that couldn't be. That's because what was the question you asked?
PD: I was wondering what happened after you went to graduate school. You went to you then went on to Bar Harbor, Maine, to work at Jackson Labs?
LL: Oh, yes. No. I went to Stanford University. They had a fellowship from Harvard, a Sheldon Fellowship, which I won. It paid \$1500. And, at the time, George Beadle, who was an associate professor at Harvard and I took a course from him decided that he wanted to go to Stanford, so he drove across the country, and I with him, because I had a fellowship at Stanford, which, for me, wasn't good. Because Beadle and I didn't get along very well. And, he was sort of a selfish guy and he was in the process of getting a divorce. So, I went with him and we stopped at his home in Nebraska, called Wahoo, Nebraska.
PD: Where he had grown up?
LL: On a farm. And we met his father and his sister, who was a nurse. Apparently, his mother died. So, he went to the University of Nebraska and then went to Cornell University. You know, he finally got a Nobel prize in physiology and medicine, which he deserved. He was a very bright person.
PD: So, you two arrived in California
LL: We arrived in California, and they had his lab all fixed for him and he started right out doing work. He worked, at that time, on Drosophila melanogaster. So, the man he had working for him got me a room in Palo Alto, Stanford. I was there a year not very happy year. So, then I came to Bar Harbor.
PD: And then who did you work with up there? And what did you do?
LL: Well, I met when I was at before we went to California, Beadle worked at Cold Spring Harbor with a man by the name Sturdevant, who was a professor at Columbia University. So, I made arrangements to drive with him, and, in the meantime, I met C.C. Little Clarence Cook Little at a lobste party in Cold Spring Harbor.
PD: In Maine? Bar Harbor?
LL: No. No. Where they went in the summer not Cold Spring Harbor, but
PD: so this was in Woods Hole that you met him that you met C.C. Little?
LL: Yes, Dr. Little. And he told me that if I needed a job at any time, to contact him. So I did that, because I wasn't happy with Beadle, and he immediately got me a fellowship to go to Bar Harbor, which I did. I drove from California to Bar Harbor. That's when I got in trouble with the Army/Air Force. They had wanted me as a private. So I found out that they had a position for Ph.D.'s called aviation physiologists. And you had to go to Randolph Field in San Antonio for six weeks and take the course there. Well, in the meantime, I got married and it wasn't long that I was on the train going to San Antonio. And my wife had the car, and she stayed around Bar Harbor a little while and then she drove home, to the middle of lowa, Marshalltown, lowa. In the meantime, we had been married.
PD: So, you were in the Army Army or Air Force?

LL: Well, at the time at that time, it was called the Army/Air Force, you know, before it was called Air Force. So, I was in the Army/Air Force for four years. After being trained, we were assigned to, you know we were in Texas and Colorado, Utah anyplace they had B-29 Air Force. That's where we were as aviation physiologist training crews to fly at 38,000 and also how to ditch their plane. It's a huge plane which wasn't too good, and a lot of them crashed. Most of the time we spent in Clovis, New Mexico, and that's where our first son, Bill, was born. And we trekked around Utah and Colorado, and Texas. It was nice. It was nice to later see green things. Okay
PD: So, after that, you came back. After you finished your military duty, didn't you return to Jackson Labs as a scientific researcher?
LL: Yes, I returned to Marshalltown, Iowa, my wife's home, because a person had offered me a job at the University of Minnesota. And I went to look at it and I didn't want it. So I came back to Marshalltown and then I went to Bar Harbor for a year, the same old job as also the scientific director, whatever that meant, because we had only about ten people in the Jackson laboratory.
PD: Had you worked had you started working on chemotherapy by this time? What was the status of your research interests?
LL: No four years in the Army/Air Force. You know, we didn't we didn't do anything in research. But I got right on joining research. The peculiar thing about Bar Harbor was nobody bothered you. They didn't tell you what to do or where to live or anything. So I was on my own, and I was interested first in carcinogenesis. And I got hold of a compound they call azo azo compound. And it was interesting because I had a summer student there whose uncle was in cancer research. I can't think of her name Rosenthal, I think. And she finally went to Johns Hopkins University and stayed in cancer research as a pathologist, which I like. And she did very well the summer that I had her. So, after that, I don't know I think I was still interested in carcinogenesis, but the fire came along and wiped out all of the mice which I had set up to send to Bethesda. And, so, there I was in Bethesda without any mice.
PD: So, when you joined the National Cancer Institute in 1947, your mouse colonies had been destroyed?
LL: Yes.
PD: How did you deal with that when you got there?
LL: Oh, I had a person by the name of Walter Heston and Howard Andervont and all of the other people worked on mice. That was the animal of choice. And they got their mice from the Jackson laboratory, which C.C. Little had worked when he was at Harvard he worked on mice and he was one of the first to inbreed mice, which means brother times sister, every generation, and has figured out when all the mice will be alike genetically. But C.C. Little was responsible for that. I don't know why. He was always loaded with bourbon [laughter]. So
PD: How did you join the National Cancer Institute in the first place? Who suggested that you apply there, or how did that come about?
LL: Well, when we moved to Bar Harbor, a very good friend, who was developed in Bar Harbor, was Walt Heston. He was a geneticist who had been prepared for the NCI. They had about ten people training in Boston, set up for the NCI, to move to Bethesda in time. And he was sent to Bar Harbor. But the other people trained in Cambridge and Boston at Harvard University. I think there were ten to twelve people that were transferred to Bethesda.
PD: So, when you got there, you recreated your mouse colony
LL: Yes.
PD: and you began working as a geneticist in the Lab of Biology with Walter Heston. Is that correct?

LL: Oh yes. Right. He was the head of the laboratory was Andervant, and Heston was under him, as a geneticist, and a very good one. And when I moved there, of course all of those people in Boston transferred to Bethesda. And also, they had a unit at the University of California, San Francisco, who did clinical work. I don't know what they did not very much. But they came so all of that was the root of the Cancer Institute the people
who were trained. And I, of course wasn't trained. I had Heston get me a fellowship. So, when I first came here, I was on a fellowship.
PD: When you arrived, what kinds of research did you do? Did you start doing chemotherapy research then? Or, how did that come about?
LL: No. I was interested in gonadectomy and leukemia, so we were interested in the in testicles and ovaries and the relationship to that's when we used the C-58 animals. I got those again from what's his name MacDowell, who was at he was in New York at Cold Spring Harbor.
PD: Okay.
LL: So, I got what I wanted from him and also other mice from various people at the NCI. And they all worked with the mice at that time.
PD: So, tell me how you got into leukemia and chemotherapy research.
LL: I don't know. I don't know except we were interested in the genetics of leukemia you know, susceptibility. And I used C-58 for that.
PD: Okay.
LL: And that was a lot of the mice that got burned were C58. I had hundreds of mice to ship. In fact, you know, Heston did all of his experiments with hundreds of mice. Today, you couldn't do it. They wouldn't have space for it [laughter]. But despite the interest that Heston had in what I was I was doing, we never did any work together.
PD: Really? I wonder why.
LL: Well, there were various reasons. One reason they moved us out of Building Six that was the Cancer Institute, and everybody set up in Building Six, which still exists. And I moved to Building Eight, which is a building next to Building 1, because they had we had space laboratory space and space for the animals. So I moved there from Building Six that separated us. At that time, though, you didn't do much collaborative work. You know, there weren't that many people and they were very good pathologists and radiologists and geneticists they were limited in what they could do, you know. So, I was surprised that, in the work that I did on development of resistance to antimetabolites, it took hundreds of mice to do that. And it was nice that we were at the National Cancer Institute.
PD: So, tell me what research was like at NCI when you first arrived what the facilities were like
LL: Very good. I was situated in the attic of Building Six. That's where they wanted me. And I had a lab there, and mostly animals. You know, you had to check the animals all the time. And everybody had was interested in cancer research. What they did was dependent upon what they were trained. As I remember our pathologists were extraordinary that good. And [Harold] "Red" Stewart had, over the years, got some young pathologists in which we didn't work with because, you know, we had to know whether we were dealing with a lymphocytic leukemia or not. And I remember this person I worked with closely Thelma Dunn. She was from Virginia and she was a very good interested in mouse leukemia. And she was the first one that classified the leukemias for Potter and for me, and also there were new young pathologists. Clyde Dawe
PD: Dawe?
LL: D-A-W-E, which we did a lot of work with. He was Ph.D., MD from Johns Hopkins and I did work with him because he was one of the first at the Cancer Institute or anywhere else that grew tumors in-vitro. You know, there were we worked with nice leukemias, but you had to transplant them from mouse to mouse. And people had to develop in-vitro. So, one of the first things I did was hire people who were interested in in-vitro development. And I remember we had a man from Wistar Institute on the fellowship and we had two people from Australia on fellowships, just to work on the development of in-vitro passage.

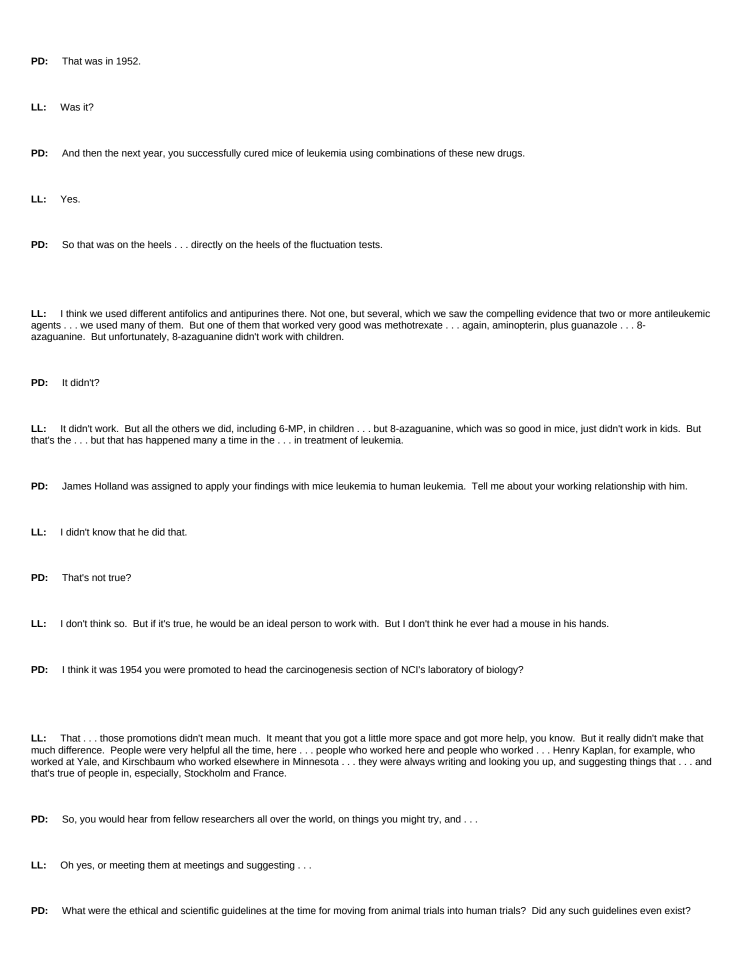
PD:	Anybody else who stands out in your mind from those early days?
at the from S fellows	Oh yes. Then, of course, with time, then you knew what people were doing and what they wanted to do, so we, besides working with people there NIH, such as virologist, Wally Rowe and the people in pathology, we got fellows in, to spend one or two years. And we had people from Australia, Sydney. We had people from Israel, Nathan Trainin, who spent two years with us. And I would say we had ten to twelve people who came as s. And one type of fellowship we liked was Eleanor Roosevelt type of and we had a man from Sydney who spent two years and a man from from the Weizmann Institute [of Science], who spent two years here.
PD:	So, you had people from all over the world coming
	We had pretty and not only that, but they really wanted me to come talk about my work in England and Stockholm and France all over the And, you know, cancer research was just beginning at that time.
PD:	What was known about it? Was there was it pretty wide open?
interes	It was wide open. And one thing that we had to develop right off was a biological system that would divert, you know? And we got one of the lantable leukemias, 1210 L-1210 that we developed from all of the in-vitro and invivo work. And that is one thing that Thelma Dunn was very sted in, classifying leukemias. And we developed it. And soon it was all over the world. And we lost our transplant. We had it by microbiological ants here in Cambridge and they lost it. So, we could always get it back, but it wouldn't be the same thing.
PD:	It was you who developed the L-1210 right?
LL:	Yes. "L" means "Law."
PD:	Did you do that by yourself, or were you collaborating?
Penns who to did a l	Well, you know, we had technicians who always helped. And we had a man by the name of Miller, who was from Millerstown State in sylvania. And we had a name by the name of Boyle who was from a little school in Hamline, Minneapolis, who was here. And then we had people book care of the mice, who developed skills. No one person removed a thymus or removed and if they so, it they were interested and they ot of work. As I I remember, Jim Miller, who helped me with the studies on resistance. He worked very hard. And I sent him to Yale University his Ph.D. because I knew Beadle did very well and he had connections with Yale. And the guy flunked out. He didn't pass his French reading exam.
PD:	So, if it was wide open if the area of cancer research was wide open, how did you hone in on this particular area?
LL:	You mean cancer research?
PD:	Chemotherapy research for leukemia.
LL:	Well, you know, the problem that we faced at one time, which Jim faced and what's his name
PD:	Dr. [Joseph] Burchenal?



LL: Jim Holland they used these compounds and they found an amazing response of the leukemic kids to 6-MP, especially one to methotrexate. And this is the first time that they cured childhood leukemia. And I think somewhere, somebody ought to reduce that in writing. You know, how they used the compounds, how many people, how many children they used and what the response was. Nobody has done that. And I would think the person who is capable of doing it is Tom Frei. I think he was the scientist of the group. So, but they Zubrod and these people and we had a laboratory of biology and I didn't have, really, too much contact with Frei and Freireich. And Holland and I became fast friends, because he liked to get lamb patties and come into our place and cook them [laughter]. He was divorced at that time. So
PD: Well, I have a question about just what a typical day in the life is like for you while you were working with these people. Or, were you pretty much in the lab and they were somewhere else? Or were you interacting constantly every day? Kind of walk me through a typical day.
LL: You mean Frei and Freireich?
PD: Frei and Freireich Dr. Holland
LL: I saw very little of them, because they were in the, you know, in the clinical part of it. The clinic had just been finished in 1953 and shortly after that, I think Zubrod got his people together. But I, you know, I knew Frei and Freireich. We lived side by side and we talked a lot. But some of those people didn't have much to do with curing leukemia, like Laszlo.
PD: He didn't?
LL: No.
PD: Well, tell me what a day was like for you then. I mean, you would get in early I mean, just, you know, when would your day begin? What would your usual schedule consist of?
LL: Well, there was a time when you didn't have much money for travel, you know. And the first time we came here, when we came here, we had to use we had to live in a house a room in a house, for eight weeks.
PD: You and your wife and your children?
LL: Yes. Well, one Bill was about two years old then. But I remember eating out at the Marriott every night. There weren't any places. And finally, they developed some apartments on Bradley Boulevard that was only for Army retirees and Army/Air Force, Navy and so forth. That was the first indication that we could get anything Bradley Boulevard.
PD: Well, tell me a little bit about the response in the scientific community to these discoveries regarding combination chemotherapy. Wasn't there a lot of resistance among other scientists to the idea that you could use a combination of drugs to cure leukemia?
LL: No, I don't think so. I don't think so. It was that people hadn't done it. You know, they had done it in bacteria in salmonella [Sal] Luria and [Max] Delbruck did it. But that was easy, you know. When you came to doing it with mice, it was very difficult because of the numbers of mice, where you kept them and took care of them, and somebody to look at the leukemias and decide what they were.
PD: You mentioned that Dr. Freireich lived next door to you?

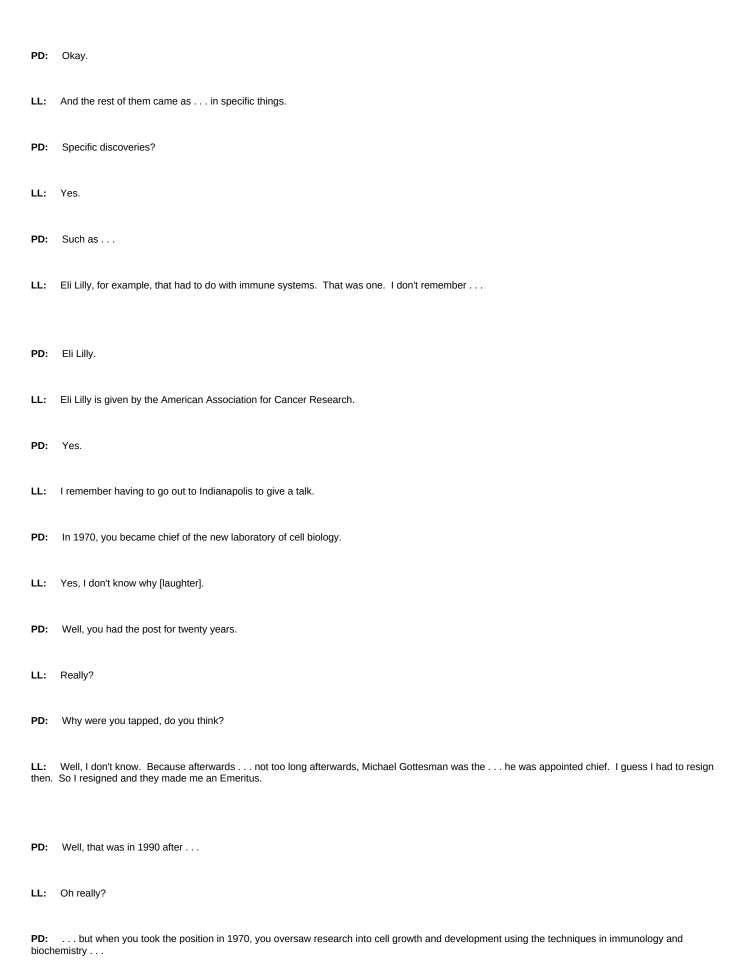
LL: Freireich, yes.

PD: finding	I read a story that it was during backyard conversations, while your children were playing, that you and he came up with the idea of trying your gs in mice on humans. Is that true?
scienti leuker	You know, that's cutting out Frei and Holland and so forth. And I think Holland was the person to push, you know, collaboration. Frei was a good ist, but Holland pushed. And they, with the new clinical facilities there, and they money that they had they had a straight shot to curing nia. I don't know what Burchenal was doing at that time. Burchenal did a lot with mice, as well as human beings. So, but, I don't think he was d, in any way, experimentally with these people here, you know.
PD:	Okay. Right. He was in New York.
LL:	Yes.
PD:	A year after you came to NCI, individual drugs started to be used for acute lymphocytic leukemia. Did your research help play a part
LL:	Oh yes.
PD:	in their being discovered?
LL:	No.
PD:	No?
LL: accura	No. I the organic chemists, Hitchings and Gertrude Elion did that on their own. What they had to tell them that, you know, biologically they are ate, I don't know. But I do remember talking with George Hitchings and he wanted always to send new drugs to treat, to test.
PD:	An experiment for which you are especially well known is the fluctuation test. Could you tell me how that came about and what it proved?
many State i didn't should for each amino amino resista	Well, we have found that some leukemic clones would respond, and some wouldn't respond, to, I think aminopterin. That's the one that took so mice and hundreds of mice which I did with this man Jim Miller, who I brought from Bar Harbor. He had a bachelor's degree from Millersville in Pennsylvania. He was very good. But we noticed that some clones clones that when you're taking off groups of cells some clones respond to methotrexate, and they found that some of the kids weren't responding to what we call aminopterin at that time. So we decided we dook at cells which are selected by drug, and which aren't selected by drug, as to what they did. So, you know, we had to use twenty or thirty mice ch little clone. And we found out that the methotrexate (aminopterin) really didn't have anything to do with it, and with the clone not responding to pterin. And when we looked closely at it, we found that true, that you had you had clones which didn't respond, that had no previous the pterin had no place in it, you know. We were using a fluctuator test which Luria and Delbruck had used for salmonella when they studied for ance to salmonella. So, it was a matter of what you selected, how many mice you used and what you were looking for. And that's why Frei, Tom ouldn't believe it, you know. But it was there.
PD: agents	So, in the fluctuation tests, were your results the first compelling evidence that combination chemotherapy might be effective where individual s would not?
LL: what v	I would think so, yes. Because we used antifolics plus antipurines and antipyrimidines. And that, especially 6-MP and methotrexate really told us vas happening.
PD:	And then not long after you conducted the fluctuation tests
LL:	That's the fluctuation test.

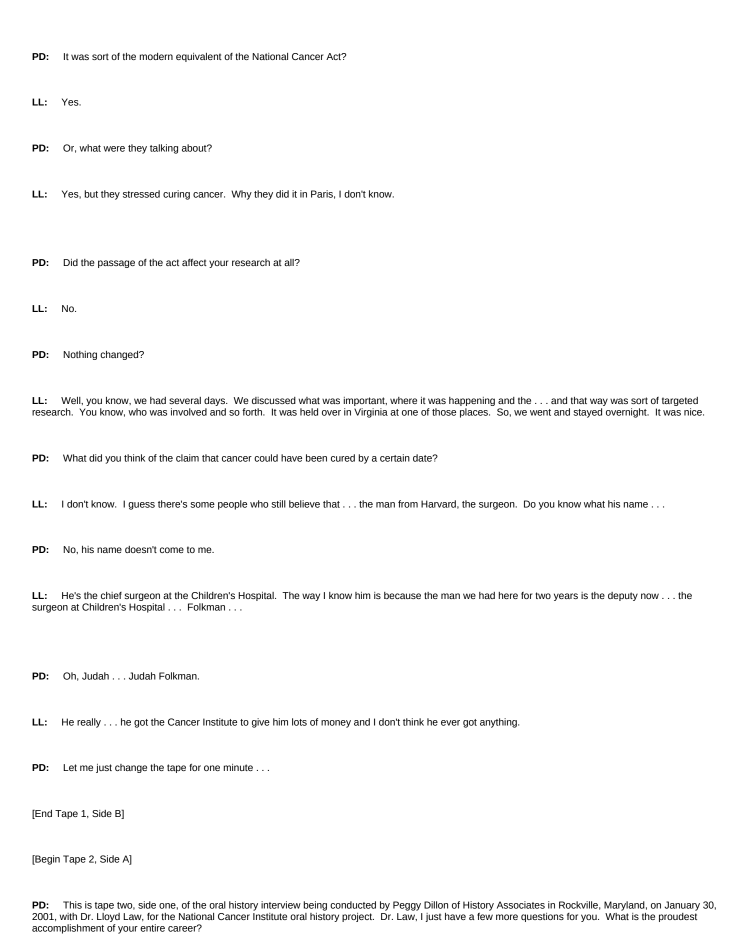


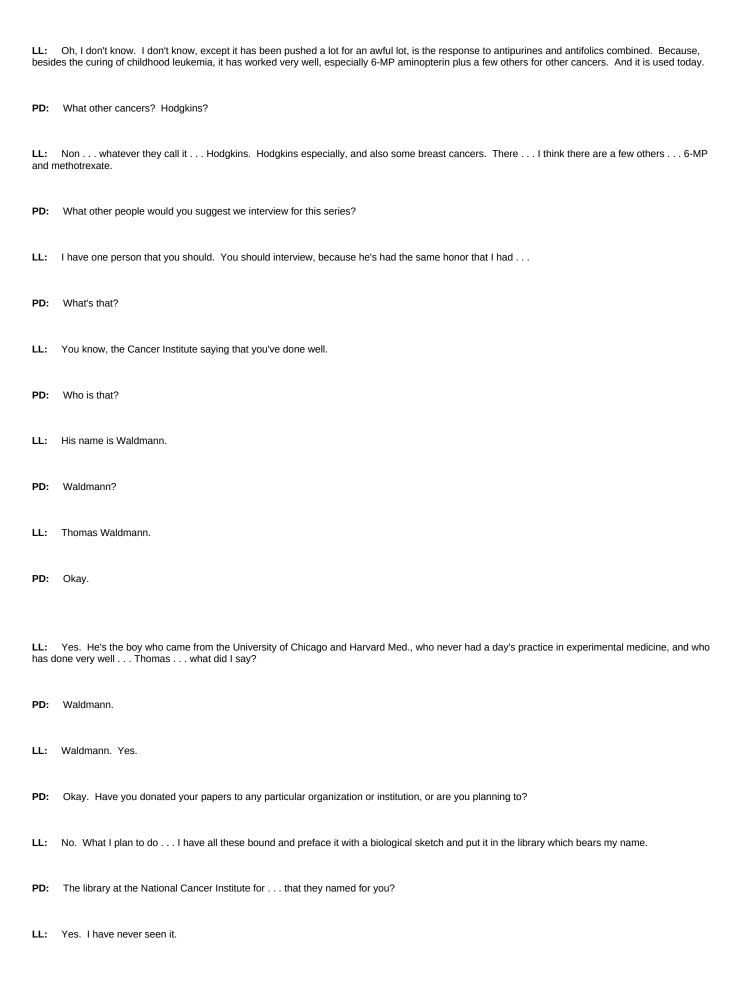
LL: them	No. I mean, the kids were dying of leukemia. And they had these six or ten clinical associates that were wanting something to do. And Zubrod put at work curing leukemia.
PD:	So how did you judge when a treatment was deemed effective enough to move it from mouse trials to human trials?
	I don't think we ever looked into that, except, you know, like 8-azaguanine, which was so good in mice it just didn't work in human beings. But I you know, it worked in mice, so you tried them in human beings. But I think you know, you said someplace, under the guidance of d. Zubrod was strictly a clinical person. And I never talked with him very much.
PD:	You were not that you didn't come into that much contact with him on a daily basis?
LL:	No, because he was in the clinical center and I was in Building Eight.
PD:	You moved buildings?
LL:	Yes, Six, Eight, the clinical center, back to Eight. It depended upon how many people we had and how much space we needed.
PD: LL:	So that's why people would go back and forth between Six and Eight? Yes.
PD:	When did you move from one to the other?
LL: you w	I don't know. [There was] very little moving. And I always got along with the people who, you know, the directors. They were always nice for what ranted.
PD:	Do any of them stand out in your mind? Ken Endicott or Carl Baker?
mone they a	Well, Carl was trained as a chemist, you know, at the University of Louisville, but he started working with mice and found that he couldn't work with So, I was close to Carl, yes, and Endicott. Endicott was trained as a pathologist, and the reason they got these other jobs was they got more y. We're thinking about people well, Thelma Dunn, who worked in the lab, stands out, and Red Stewart, another pathologist, and Clyde Dawe all were interested in what happened in the lab, and they didn't really have much to do with human pathology. You know, that's what they wanted to so, it's been a good place to work, you know.
PD:	You studied a couple of different areas
LL:	Yes, the area that we are discussing was the first area
PD:	Yes. Tell me about how you moved into other research interests.

with, v time o cell m	Well, because of people, you know and what was happening. People were interested in viruses polyoma virus, which we did a lot of work with Wally Rowe and then we were interested, for a long time, in cell antigens which have to do with immune system, which we spent an awful lot of n, in mice. Because at that time, we had developed in vitro methods and we had also a person from Stanford University who was interested in H-2 arkers, because he spent two years with us and trying to characterize those normal antigens. And he went back to Stanford. His name was Strober. But he did a lot of good work.
PD:	Well, on the subject of virus oncogenicity, what would you say your major contributions were in that area?
	Well, in conjunction with Wally Rowe and Clyde Dawe, we defined the conditions necessary for production of tumors by polyoma virus and also by ukemia viruses, which we found were effective. So, it was what they wanted to do and what we wanted to do together. It made things sting. And all these people were wanting to collaborate. They sent Heston and Andervont and I never got together.
PD:	The three of you never worked together?
LL:	No.
PD:	How about your work on immune mechanism cancer induction and immune mechanisms?
the moback ι	Well, we used some tumors that were induced by the chemical called methylcholanthrene or 910-Dimethyl, which developed in mice and we had several of them were immune to just antigens that you prepared by solubilizing. So we did a lot of work for that, and also with Vincent Hearing on elanomas recently we did in attempting to characterize these antigens which we got by the usual methods of solubilizing antigens. And that all turns upon the man Strober who came to us from Stanford University, who was interested in the H-2 system of antigens, which he solubilized. So that d out very well his two years. That was the beginning, I think, of when we used sarcomas, melanomas in attempts to in attempt to get active al.
PD:	To get what kind of material?
LL:	Active material.
PD:	Oh, okay.
LL:	Yes. Most of the honors that I have for that, are for immune system. The first one was especially to resistance.
PD:	Which honor?
LL:	Rosenthal.
PD:	The Rosenthal Award?
LL:	Yes. That was early.
PD:	Okay. The Anne Frankel Rosenthal Award, in 1955. What was the particular discovery that, for which you won that award?
LL:	The fluctuation tests.



LL:	I guess that's when we started looking at the antigens, which would be solubilized there in 1970, yes.
PD:	Was technology changing at that time? Was molecular genetics coming on to the scene?
LL: then,	No. Molecular genetics had nothing to do with it. But, you know, you would always have things that happened. I don't remember what happened but it was at a time when immunology played a good role, you know.
PD:	What would you say were the positive aspects of being at NCI during your many years there? What were the highlights of your career?
LL: difficu	People. People who did things, and discussed them. There were good scientists. The NCI tried to get the best that they could get, and it was verill to get a job a permanent job at the NCI or NIH, but particularly NCI. And also the NCI had more money than any other institutes.
PD:	Even from the start?
LL:	Even from the start. You know, a person who had quite a lot to be in and quite a lot to do in developing the NCI was Clarence Cook Little.
PD:	Did he?
LL:	Yes. He was very active in developing the NCI.
PD:	Any other particularly positive aspects of your time there?
LL: collab	porate with you, and a good place to work, you know. I finally moved to Building Thirty Seven, which was a wonderful place for the laboratory of cell
PD:	Not long after you became head of the lab of cell biology, the National Cancer Act was passed. Were you present at the signing of that?
LL:	Yes.
PD:	You were there?
LL:	Yes.
PD:	What was that like?
	Oh, a lot of interesting people. We had a meeting and then we signed. That was it. We had a discussion for several days of what we were doing what we expect. And I remember meeting people I had seen before at this meeting. They had a meeting like that last year in France. They invited body back to sign something, that they would spend all their time and money on cancer research.





PD:	Okay.
LL: biogra	But the, I invite Gottesman, invite Hearing and invite Appella to do the same thing then they have all the reprints and with the phical sketch
PD:	That's great. What are you ties to the National Cancer Institute right now? Aren't you an Emeritus investigator?
LL:	Yes. That means they give you a little money and they give you a place to work and also a place to park and they're good
PD:	Do you go in much?
LL:	No. I used to go in about two or three days a week, but recently, with my legs, I have difficulty walking. So maybe I'll get that fixed up.
PD:	Any final thoughts that you'd like to share that I haven't thought to ask you?
LL:	You can come back and ask. I'll think about it.
PD:	Okay. We'll do that. Well, thank you so much for your time today.
remer	You know, it couldn't be better, what has happened as far as my life scientifically is now, because everything has turned out wonderfully. And I just niber today of going through a huge apartment in Paris by this French scientist named Lacassagna, who I got to know very well. But he invited me omebody else to have the best roast lamb I ever had, in his apartment in the middle of Paris. And there are so many things that happened that way, now?
PD:	It was a very good career?
LL:	Yes. But I'll think of the special things.
PD:	All right. We can revisit them another time. Thank you very much.
LL:	You can come back.
[End o	of Interview]
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